Security Classification of This Report Has Been Cancelled

COPY NO. 124

NACA RM No. E8117a

6571

E8I





# RESEARCH MEMORANDUM

THEORETICAL PERFORMANCE OF DIBORANE AS A ROCKET FUEL

By Vearl N. Huff, Clyde S. Calvert and Virginia C. Erdmann

Lewis Flight Propulsion Laboratory Cleveland, Ohio



CLASSIFIED DOCUMENT

The contains classified information affecting to Defense of the United States with USC 50:51 transmission or the revelation of its unauthorised pers. It is not present the control of the Control of the Control of the Control of the United States of the United S

NATIONAL ADVISORY COMMITTEE FOR AERONAUTICS

WASHINGTON January 10, 1949

319.98/13

Security Classification of This Report Has Been Cancelled

NACA RM No. E8117a



#### NATIONAL ADVISORY COMMITTEE FOR AERONAUTICS

#### RESEARCH MEMORANDUM

THEORETICAL PERFORMANCE OF DIBORANE AS A ROCKET FUEL

By Vearl N. Huff, Clyde S. Calvert and Virginia C. Erdmann

#### SUMMARY

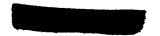
Theoretical performance data based on equilibrium isentropic expansion and constant-composition (frozen) isentropic expansion from a combustion-chamber pressure of 20.4 atmospheres (300 lb/sq in. absolute) to an ambient pressure of 1 atmosphere are presented for a range of mixtures for four rocket-propellant combinations of diborane with liquid fluorine, liquid fluorine exide, liquid exygen, and 100-percent hydrogen perexide. The theoretical data include combustion-chamber and nozzle-exit temperatures, specific impulse, and volume specific impulse. Composition and mean molecular weight of the reaction products are given for both the combustion chamber and the nozzle exit.

The maximum specific impulse for all the combinations occurred in the fuel-rich region. On the basis of maximum specific impulse, the four oxidants reacting with diborane assumed the following order: liquid fluorine, liquid fluorine oxide, liquid oxygen, and 100-percent hydrogen peroxide. On the basis of calculated maximum volume specific impulse, the order of the four oxidants reacting with diborane was: liquid fluorine oxide, liquid fluorine, 100-percent hydrogen peroxide, and liquid oxygen.

### INTRODUCTION

Considerable interest has been shown in some of the boron compounds as rocket propellants because their high heat of combustion per unit weight indicates a high specific impulse.

Pentaborane and diborane are two of the more significant of the boron hydrides under consideration because chemically they are sufficiently stable to permit synthesis and use. Pentaborane has several advantages over diborane including higher density and boiling point, but diborane is more readily available and more data on its heat of combustion exist.



2

Computed performance of diborane and liquid oxygen for three mixture ratios under equilibrium-expansion conditions and for a range of mixture ratios under frozen-expansion conditions is presented in reference 1. A theoretical and experimental investigation of boron compounds as rocket fuels is being conducted at the NACA Lewis laboratory and calculations are reported herein that were made from December 1947 to May 1948. These computations extend the mixture range of reference 1 for diborane and liquid oxygen and, in addition, give performance over a considerable mixture range of diborane with liquid fluorine, liquid fluorine oxide, and hydrogen peroxide. The results include combustion-chamber temperature, nozzle-exit temperature, specific impulse, volume specific impulse, composition, and mean molecular weight as functions of mixture ratio.

#### METHOD OF CALCULATION

The calculation of specific impulse involves the determination of the gas composition and the temperature in the combustion chamber and at the nozzle exit. The products of reaction were assumed to expand from a combustion-chamber pressure of 20.4 atmospheres (300 lb/sq in. absolute) to an ambient pressure of 1 atmosphere. The ideal gas laws were used.

The molecules considered to be present in the gas phase for the appropriate reactions were: atomic hydrogen H, hydrogen  $H_2$ , water vapor  $H_2$ 0, hydroxyl radical OH, hydrogen fluoride HF, boron hydride BH, atomic oxygen O, oxygen O2, boron trioxide  $H_2$ 0, boron oxide B0, atomic fluorine F, fluorine F2, boron trifluoride  $H_2$ 1, boron fluoride BF, atomic boron B, and diatomic boron  $H_2$ 2. At nozzle-exit temperatures below  $H_2$ 2000 K, liquid boron trioxide  $H_2$ 203 was also included but boron B in the liquid and solid states, which could also be present, was neglected. The boiling point of boron is given as  $H_2$ 2823 K in reference 2 (p. 1750). For many of the calculations, the liquid and solid states of boron would not be present and, for the remaining calculations, the effect on specific impulse is probably small.

The theoretical performance was calculated on the basis of constant enthalpy from the fuel and the oxidant at the assumed initial state of the propellants (tank conditions) to the state of the reaction products in the combustion chamber. The term enthalpy  $H_T^{O}$  is defined by



$$\mathbf{H}_{\mathbf{T}}^{\mathbf{O}} = \int_{0}^{\mathbf{T}} \mathbf{c}_{\mathbf{p}} d\mathbf{T} + \mathbf{H}_{0}^{\mathbf{O}}$$

where

HO heat of formation at 0° K, calories per mole

T temperature, OK

cp specific heat at constant pressure, calories per mole per CK

The superscript o denotes the thermodynamic standard reference state of unit activity; the subscript T denotes the absolute temperature in degrees Kelvin. After the enthalpy of the fuel plus the oxidant at initial states was calculated, the combustion temperature was obtained by simultaneously solving the equations for equilibrium, mass balance, and enthalpy by a method of successive approximations. The equation for enthalpy of the reaction is

$$\mathbf{H}_{\mathbf{T}}^{\mathsf{O}}$$
 (reactants)=  $\sum_{i} \mathbf{n}_{i}$  ( $\mathbf{H}_{\mathbf{T}}^{\mathsf{O}}$ )<sub>1</sub> (products of reaction)

where

 ${\tt H}_{
m T}^{
m O}$  (reactants) enthalpy of reactants at initial states, calories

n<sub>i</sub> moles of product i

 $(\mathbb{F}_{\mathrm{T}}^{\mathsf{O}})_{1}$  enthalpy of product i, calories per mole

The nozzle-exit temperature was calculated on the assumption that chemical equilibrium prevailed throughout expansion (equilibrium expansion) and on the assumption that no chemical recombination took place (frozen expansion). In each case, isentropic expansion was assured by comparing the entropy of the fluid at the exit S<sub>e</sub> to the entropy of the fluid in the combustion chamber S<sub>c</sub>.

$$S_{c} = S_{e} = \frac{1}{\sum_{i} n_{i} M_{i}} \left[ \sum_{i} n_{i} \left( S_{T}^{o} \right)_{i} - R \sum_{i} n_{i} \log_{e} p_{i} \right]$$



NACA RM No. E8117a

where

4

M, molecular weight of product i

 $(S_{\mathrm{T}}^{\circ})_{,}$  absolute entropy of product 1, calories per mole per  ${}^{\circ}K$ 

R gas constant, 1.98714 calories per mole per <sup>C</sup>K

p, partial pressure of product i, atmospheres

Specific impulse I (lb-sec/lb) was calculated from the difference in enthalpy between the combustion chamber and the nozzle exit by the equation

I = 9.328 
$$\sqrt{\frac{\frac{H_T^o}{\sum n_1 M_1}}{\sum n_1 M_1}} - \left(\frac{\frac{H_T^o}{\sum n_1 M_1}}{\sum n_1 M_1}\right)_e$$

Volume specific impulse  $I_d$  (lb-sec/(cu ft)(62.43) or gram-sec/cc) is also included.

#### THERMOCHEMICAL DATA

The heats of formation of diborane  $B_2B_6$ , boron trioxide  $B_2O_3$ , and boron trifluoride BF3 were taken from reference 3. Some uncertainty exists concerning the thermochemical data for the various states of boron and its compounds. The heat of combustion of diborane with oxygen, however, is not subject to so much question and is given by reference 3 as 510 kilocalories per mole. The value of the heat of formation of B2H6 has been given as -44 kilocalories per mole in reference 3, -29.5 kilocalories per mole in reference 4, and recently a value of 26 kilocalories per mole in reference 5. (Heat liberated is considered to be negative.) All the data presented herein are based on a heat of formation of -44 kilocalories per mole for diborane. One calculation, however, was made for the stoichiometric mixture of diborane and liquid oxygen using the value of 26 kilocalories per mole and the specific impulse was 2.5 percent lower than by using the value of -44 kilocalories. The heat of transition of  $B_2O_3$  (amorphous) to  $B_2O_3$  (crystalline) was neglected. The effect of this assumption is small and will change the specificimpulse values on the order of 0.7 percent.



520T



The heats of formation of hydrogen fluoride HF and atomic fluorine F were taken from reference 6. The lower value ((1/2)(63.5 k-cal)) given for F was used because it is the basis of the thermodynamic properties of F and  $F_2$  (reference 6). The heats of formation of  $H_2O$ , O, OH, and H were obtained from reference 7. The sensible enthalpies and entropies of HF, F, and  $F_2$  were taken from reference 6 and those of  $H_2O$ , O,  $O_2$ , OH, H, and  $H_2$  from reference 7.

The heats of fusion and vaporization of B and B<sub>2</sub>O<sub>3</sub>, the heats of formation of BO and B, and the sensible enthalpies and free energies of B<sub>2</sub>O<sub>3</sub>, B, B<sub>2</sub>, and BO to 5000° K were taken from reference 8. The functions for B were extended to 6000° K in the same manner in which the calculations were made in reference 8. Free energy, sensible enthalpy, and entropy of BF<sub>3</sub> in the ideal gas state from 298.16° to 1000° K were taken from reference 9 and extended to 6000° K from spectroscopic data of reference 10, by assuming that the BF<sub>3</sub> molecule is a rigid rotator and a harmonic oscillator.

Similar calculations were also made for BH and BF from the spectroscopic data of reference 11. The  $^{3}\pi$  state was considered the ground state for BF because no other information was found in the literature. The values of enthalpy, entropy, and free energy for BF3, BF, BH, and B are listed in tables I, II, and III, respectively.

The propellants were taken as liquids at the following initial temperatures:

Propellant (100 percent)	Initial temperation (CK)				
Diborane	298.16				
Fluorine	85.16				
Fluorine oxide	128.30				
Oxygen	90.10				
Hydrogen peroxide	298.16				

Liquid  $B_2H_6$  was used at 298.16  $^{\rm O}K$  because no specific-heat data were available to adjust the initial temperature to or below the boiling point. The effect on specific impulse of using this



NACA RM No. E8117a

6

initial temperature for  $B_2H_6$  is small and an error in the opposite direction and of the same order of magnitude was introduced when the heat of transition of  $B_2O_3$  (amorphous) to  $B_2O_3$  (crystalline) was neglected.

The density of diborane increases as the temperature is lowered and therefore the highest density reported in reference 12 (0.4818 gram/cc at -129°C) was used for the computation of volume specific impulse. Additional physical and thermochemical properties of diborane and the oxidants considered were taken from references 13 to 16 and are given in table IV.

#### RESULTS AND DISCUSSION

The performance parameters of diborane reacting with liquid fluorine, liquid fluorine oxide, liquid oxygen, and loo-percent hydrogen peroxide are plotted against the percent by weight of fuel in the mixture in figure 1. The quantities plotted for each combination for both frozen and equilibrium expansions are combustion-chamber temperature  $T_{\rm c}$ , nozzle-exit temperature  $T_{\rm e}$ , specific impulse I, volume specific impulse Id, and mean molecular weight in combustion chamber  $M_{\rm c}$ . The mean molecular weight at the nozzle exit  $M_{\rm e}$  was also included for equilibrium expansion.

The curves of combustion-chamber temperature for the combinations considered reached a maximum near the stoichicmetric mixture for all of the cases except fluorine oxide, which did not reach a maximum in the range considered. The highest combustion-chamber temperature obtained was 5380° K for the fluorine-diborane reaction (figs. 1(a) and (b)). The hydrogen peroxide (figs. 1(g) and (h)) and oxygen (figs. 1(e) and (f)) combinations gave temperatures substantially lower than the fluorine. The combustion-chamber temperatures reached maximum computed values of 4750°, 4022°, and 3230° K for fluorine oxide, oxygen, and hydrogen peroxide, respectively. The nozzle-exit-temperature curves for both the frozen and the equilibrium expansions follow a trend similar to those of the combustion-chamber temperature and reach a maximum in the region of the stoichicmetric mixture.

The specific-impulse curves for all the combinations considered reached maximum values in the fuel-rich region as a result of the liberation of free hydrogen, which reduces the average molecular weight of the products of combustion. The trends of the specific-impulse curves based on both frozen and equilibrium expansions are similar. The maximum deviation between the equilibrium-expansion and the frozen-expansion specific-impulse values was 8.08 percent of the equilibrium values.

NACA RM No. ESIl7a



7

The maximum specific-impulse values that were calculated for equilibrium expansion are listed in the following table, together with the corresponding volume specific impulse.

Oxidant	Fuel (percent by wt)	Maximum specific impulse, I (lb-sec/lb)	Volume specific impulse, I <sub>d</sub> lb-sec (cu ft)(62.43)	Combustion- chamber temperature, (°K)	Tc
Fluorine Fluorine oxide	13.63 20.41	·322.4 316.2	309.8 375.3	5240 4460	
Oxygen Hydrogen peroxide	36.58 35.18	311.4 289.0	236.7 245.3	37 <b>4</b> 0 2850	

Because the densities of liquids F2, F20, O2, and H2O2 are greater than that of liquid B2H6, the maximum volume specific impulse occurred at a mixture ratio less fuel rich than that required for maximum specific impulse. The general trend of the volume-specific-impulse curves is similar for both the equilibrium and frozen expansions for all the fuel-oxidant combinations. For the fluorine compounds, the maximum volume specific impulse occurs at a lower percentage of fuel than was considered. The maximum volume-specific-impulse values that were calculated for equilibrium expansion are listed in the following table, together with the corresponding specific impulse values.

Oxidant	Fuel (percent by wt)	Maximum volume specific impulse, Id lb-sec (cu ft)(62.43)	Specific impulse, I (lb-sec/lb)	Combustion- chamber temperature, (°K)	T <sub>c</sub>
Fluorine oxide	12.030	424.2	302.3	<b>4</b> 750	
Fluorine	9.853	317.0	315.5	5380	
Hydrogen peroxide	11.950	300.9	259.0	3218	
Oxygen	25.710	243.2	288.2 .	4022	



8

NACA RM No. ESI17a

The mean molecular weight at both the combustion chamber and the exit decreased as the percent by weight of fuel increased. In the case of the  $\rm H_2O_2$  reaction with greater than 35 percent fuel, liquid  $\rm B_2O_3$  appeared at the exit condition and caused the molecular weight to increase. For all propellant combinations considered, the mean molecular weight was from 12.0 to 1.7 percent higher at the exit than in the combustion chamber.

The calculated values of the various performance parameters for each of the propellant combinations and for several mixture ratios are listed in table V.

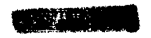
The compositions of the products of combustion in the combustion chamber and at the nozzle exit are shown in figure 2. The mole fraction of each product present in the gas mixture for each of the combinations is plotted against the percent by weight of fuel.

The curves of composition are similar for the combustion-chamber and nozzle-exit conditions for any one propellant combination (fig. 2). The general trend of a given molecule is the same for all propellant combinations. As the mixture becomes more fuel rich, B2, B, and H2 increase and F2, F, O2, and O decrease. The molecules H2O and HF are principal constituents throughout most of the mixture range considered, but are reduced by boron at the extreme fuel-rich region. The molecules BF3 and B2O3 are comparatively constant throughout the range of mixture ratios considered. Certain molecules reach a maximum (for example, H) or a minimum (for example, BF3) because of the opposing effects of the variation of temperature and mixture ratio.

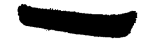
#### SUMMARY OF RESULTS

Theoretical performance data, which were based on both frozen and equilibrium expansions, obtained over a range of mixtures for the four rocket propellant combinations of diborane with liquid fluorine, liquid fluorine oxide, liquid oxygen, and 100-percent hydrogen peroxide at a reaction pressure of 20.4 atmospheres (300 lb/sq in. absolute) and a nozzle-exit pressure of l atmosphere, are summarized as follows:

1. The maximum calculated specific-impulse values in pound-seconds per pound for diborane reacting with liquid fluorine, liquid fluorine oxide, liquid oxygen, and 100-percent hydrogen peroxide, were 322.4, 316.2, 311.4, and 289.0, respectively, for equilibrium expansion.



NACA RM No. E8Il7a



9

- 2. The maximum specific impulse for all the propellant combinations occurred in the fuel-rich region at a combustion-chamber temperature less than maximum.
- 3. The specific impulse for equilibrium expansion was a maximum of 8.08 percent greater than that for frozen expansion.
- 4. The maximum calculated volume-specific-impulse values (lb-sec/(cu ft)(62.43)) for the mixture range considered were 424.2, 317.0, 300.9, and 243.2 for liquid fluorine oxide, liquid fluorine, l00-percent hydrogen peroxide, and liquid oxygen, respectively, for equilibrium expansion.

Lewis Flight Propulsion Laboratory,
National Advisory Committee for Aeronautics,
Cleveland, Ohio.

#### REFERENCES

- 1. Anon.: A Compilation of Computed Specific Impulse Values. Project RAND, RA 15049, Battelle Memorial Inst., Sept. 2, 1947. (Subcontract under AAF Contract No. W33-038 ac-14105 to Douglas Aircraft Co., Inc.)
- 2. Anon.: Handbook of Chemistry and Physics. Charles D. Hodgman, ed., Chem. Rubber Pub. Co. (Cleveland), 29th ed., 1945.
- 3. Roth, W. A., und Borger, Erika: Zur Thermochemie des Bors. Berichte d. D. Chemischen Gesellschaft, Jahrg. 70, Nr. 2, Jan. 6, 1937, S. 48-54.
- 4. Roth, W. A.: The Thermochemistry of Boron. Chem. Abs., vol. 41, no. 17, Sept. 10, 1947, column 5373f. (Abs. from Z. Naturforsch, Vol. 1, 1946, pp. 574-576.)
- 5. Eggersgluess, W., Mazurkiewicz, A., and Parker, W. G.: The Heat of Formation of Boron Trioxide. Rep. No. CHEM. 433, R.A.E., July 1947.
- 6. Kiehl, S. J., Jr., and Moore, J. R.: Propellants for Supersonic Vehicles: Liquid Fluorine. Project RAND, RA 15407, Battelle Memorial Inst., Aug. 12, 1947. (Subcontract under AAF Contract No. W33-038-ac-14105 to Douglas Aircraft Co., Inc.)





- 7. Hirschfelder, J. O., McClure, F. T., Curtiss, C. F., and Osborne, D. W.: Thermodynamic Properties of Propellant Gases. NDRC Rep. No. A-116, Nov. 23, 1942. (Abs. Bib. Sci. Ind. Reps., vol. 2, no. 10. Sept. 6, 1946, p. 743, PB 28531.)
- 8. Wacker, Paul F., Wooley, Harold W., and Fair, Myron F.: Thermodynamic Properties and Gaseous Equilibria of Boron, Oxygen and the Oxides of Boron. Tech. Rep., Heat and Power Div., Nat. Bur. Standards, Jan. 25, 1945. (Bur. Aero., Navy Dept.)
- 9. Spencer, Hugh M.: Thermodynamic Properties of Gaseous Boron Trifluoride, Boron Trichloride, and Boron Tribromide. Jour. Chem. Phys., vol. 14, no. 12, Dec. 1946, pp. 729-732.
- 10. Herzberg, Gerhard: Infrared and Raman Spectra of Polyatomic Molecules. D. Van Nostrand Co., Inc. (New York), 1945.
- 11. Herzberg, Gerhard: Molecular Spectra and Molecular Structure.
  I. Diatomic Molecules, ch. VIII. Prentice-Hall, Inc. (New York),
  1939, p. 484.
- 12. Laubengayer, A. W., Ferguson, R. P., and Newkirk, A. E.: The Densities, Surface Tensions and Parachors of Diborane, Boron Triethyl and Boron Tribromide. The Atomic Parachor of Boron. Jour. Am. Chem. Soc., vol. 63, no. 2, Feb. 1941, pp. 559-561.
- 13. Maass, O., and Hatcher, W. H.: The Properties of Pure Peroxide.
  I. Jour. Am. Chem. Soc., vol. XLII, July-Dec. 1920, pp. 2538-2569.
- 14. Anon.: Tables of Selected Values of Chemical Thermodynamic Properties. Nat. Bur. Standards, Dec. 31, 1947.
- 15. Bichowsky, F. Russell, and Rossini, Frederick D.: The Thermochemistry of the Chemical Substances. Reinhold Pub. Corp. (New York), 1936.
- 16. Anon.: International Critical Tables. Vol. 3. McGraw-Hill Book Co., Inc., 1928, pp. 20, 203, 214; vol. 7, 1930, pp.11, 212.



TABLE I - ENTHALPY OF GASES

TABLE I - ENTHALPI OF GASES							
		Enthalpy, H <sub>T</sub>		mole			
Temperature ( <sup>O</sup> K)	BF3	BF	BH	В			
1000	14,007	7,617	7,188	4,997			
1100	15,833	8,465	7,972	5,494			
1200	17,683	9,319	8,767	5,991			
1300 1400	19,551	10,179	9,574	6,487			
1500	21,435 23,332	11,043 11,911	10,387 11,213	6,984 7,481			
1600	25,239	12,782	12,043	7,978			
1700	27,154	13,656	12,880	8,475			
1800	29,077	14,532	13,721	8,971			
1900	31,007	15,409	14,568	9,468			
2000	32,942	16,288	15,419	9,965			
2200 2100	34,882 36,825	17,169 18,050	16,273	10,462			
2300	38,773	18,933	17,130 17,990	10,959 11,455			
2400	40,723	19,817	18,853	11,952			
2500	42,677	20,701	19,718	12,449			
2600	44,632	21,587	20,586	12,946			
2700	46,591	22,473	21,455	13,443			
2800	48,551	23,359	22,325	13,939			
2900 3000	50,512	24,246	23,197	14,436			
3100	52,476 54,440	25,133 26,021	24,071 24,946	14,933 15,430			
3200	56,407	26,910	25,822	15,927			
3300	58,374	27,798	26,699	16,423			
3400	60,342	28,687	27,577	16,920			
3500	62,312	29,576	28,456	17,417			
3600	64,283	30,466	29,335	17,914			
3700 3800	66,254 68,226	31,356 32,246	30,216 31,097	18,411 18,907			
3900	70,198	33,136	31,979	19,404			
4000	72,172	34,026	32,861	19,901			
4100	74,146	34,917	33,744	20,398			
4200	76,121	35,808	34,627	20,895			
4300	78,097	36,699	35,511	21,391			
4400 4500	80,072 82,045	37,590 38,481	36,396 37,280	21,888			
4600	84,025	39,372	37,280 38,166	22,385 22,882			
4700	86,001	40,264	39,051	23,379			
4800	87,977	41,155	39.937	23,875			
4900	89,957	42,047	40,823	24,372			
5000 5100	91,935	42,939	41,710	24,869			
5200	93,913 95,893	43,831 44,723	42,597 43,484	25,366 25,863			
5300	97,872	45,615	44,371	25,863 26,360			
5400	99,851	46,507	45,259	26,857			
5500	101,831	47,399	46,147	27,354			
5600	103,810	48,291	47,035	27,851			
5700	105,792	49,184	47,923	28,349			
5800 5900	107,771	50,076	48,812	28,845			
6000	109,751	50,968 51,861	49,700	29,343			
	111,736	51,861	50,589	29,840			





TABLE II - ENTROPY OF GASES AT 1 ATMOSPHERE

		Entropy, S <sub>T</sub> , cal/mole -							
Temperature (°K)	BF <sub>3</sub>	BF	BH	В					
1000	79.293	61.764	40 703	40,660					
1100	81.033	62.572	49.703	42.662					
1200	82.642	63.315	50.450	43.112					
1300	84.138		51.143	43.538					
1400	85.534	64.003	51.792	43.941					
1500		64.644	52.392	44.320					
1600	86.842	65.243	52.960	44.676					
1700	88.073	65.805	53.496	45.009					
1800	89.234	66.334	54.003	45.318					
1900	90.333	66.835	54.484	45.604					
2000	91.377	67.309	54.942	45.866					
	92.369	67.760	55.378	46.105					
2100	93.316	68.190	55.795	46.342					
2200	94.220	68.600	56.194	46.573					
2300	95.085	68.993	56.577	46.795					
2400	95.915	69.369	56.944	47.008					
2500	96.713	69.730	57.297	47.214					
2600	97.480	70.077	57.637	47.411					
2700	98.219	70.411	57.965	47.600					
2800	98.939	70.734	58.287	47.781					
2900	99.620	71.045	58.588	47.955					
3000	100.286	71.346	58.884	48.119					
3100	100.930	71.637	59.171	48.276					
3200	101.554	71.919	59.449	48.431					
3300 3400	102.160	72.192	59.719	48.582					
3500	102.747	72.458	59.981	48.729					
3600	103.318	72.715	60.236	48.875					
3700	103.873	72.966	60.483	49.015					
3800	104.413	73.210	60.725	49.154					
3900	104.940	73.447	60.959	49.288					
4000	105.452	73.678	61.189	49.421					
4100	105.951	73.904	61.412	49.549					
4200	106.439 106.915	74.124	61.630	49.675					
4300	106.915	74.338	61.843	49.796					
4400	107.834	74.548	62.051	49.916					
4500	108.278	74.753 74.953	62.254	50.031					
4600	108.712	75.149	62.450	50.143					
4700	109.137	75.341	62.648	50.252					
4800	109.553	75.529	62.838	50.359					
4900	109.962	75.712	63.025	50.461					
5000	110.361	75.712	63.207 63.386	50.561 50.657					
5100	110.753	76.069	63.562	50.657					
5200	111.138	76.243	63.734	50.750					
5300	111.514	76.412	63.903	50.652					
5400	111.884	76.579	64.069	51.040					
5500	112.248	76.743	64.232	51.131					
5600	112.604	76.903	64.392	51.221					
5700	112.955	77.062	64.549	51.309					
5800	113.299	77.217	64.704	51.395					
5900	113.637	77.369	64.856	51.480					
6000	113.972	77.519	65.005	51.564					
			001000	074007					



NACA



TABLE III - FREE-ENERGY FUNCTION OF GASES

	Free-e	nergy function		T - HO	
		cal/mole	• <b>-</b> °K		
Temperature (°K)	BF <sub>3</sub>	BF	ВН	В	
1000	65.286	54.147	42.515	37,665	
1100	66.639	54.876	43.203	38.115	
1200 1300	67.907 69.098	55.549	43.836	38.544	
1400	70.223	56.173 56.756	44.428	38.949	
1500	71.288	57.302	44.971 45.485	39.329	
1600	72.299	57.816	45.969	39.689 40.021	
1700	73.261	58.302	46.427	40.331	
1800	74.179	58.762	46.861	40.619	
1900 2000	75.057	59.199	47,275	40.881	
2100	75.898 76.705	59.616 60.014	47.669	41.123	
2200	77.481	60.395	48.046 48.408	41.358	
2300	78.228	60.761	48.755	41.592 41.814	
2400	78.947	61.111	49.088	42.026	
2500	79.642	61.449	49.409	42.234	
2600 2700	80.314	61.774	49.719	42.431	
2800	80.963 81.600	62.088	50.019	42.621	
2900	82.202	62.391 62.684	50.313	42.802	
3000	82.794	62.968	50.589 50.860	42.977 43.142	
3100	83,369	63.243	51.134	43.297	
3200	83.927	63.510	51.379	43.454	
3300 3400	84.470	63.768	51.628	43.606	
3500	84.999 85.515	64.020	51.870	43.753	
3600	86.017	64.265 64.503	52.105	43.899	
3700	86.507	64.735	52.335 52.558	44.038	
3800	86.985	64.961	52.776	44.178 44.314	
3900	87.452	65,182	52.989	44.446	
4000	87.908	65.397	53.197	44.574	
4100 4200	88.354	65.607	53.400	44.700	
4300	88.791 89.218	65.813 66.013	53.598	44.820	
4400	89.636	66.210	53.792 53.982	44.941	
4500	90.046	66.402	54.166	45.056 45.169	
4600	90.446	66.590	54.350	45.278	
4700 4800	90.839	66.774	54.529	45.384	
4900	91.224	66.954	54.704	45.487	
5000	91.603 91.974	67.131 67.305	54.876 55.044	45.587	
5100	92.339	67.475	55.044 55.210	45.683 45.782	
5200	92.697	67.642	55.372	45.879	
5300	93.048	67.806	55.531	45.973	
5400 5500	93.393	67.967	55.688	46.066	
5500 5600	93.733 94.067	68.125	55.842	46.158	
5700	94.395	68.280 68.433	55.993	46.247	
5800	94.718	68,583	56.142 56.288	43.335 46.422	
5900	95,036	68.730	56.432	46.507	
6000	95.349	68.876	56.574	46.590	

102

## TABLE IV - PHYSICAL-CHEMICAL PROPERTIES OF PROPELLANTS

Temperatures in superscripts, °C. References in parentheses.

						os m pe			
Propellants	Mole- cular weight M	Density (gram/cc)	Enthalpy of formation, $\Delta H_f$ (k-cal/mole)	Enthalpy of vapor- ization, AH (k-cal/mole)	Boil- ing point (°C)	Freez- ing point (°C)	Vapor pressure (mm)	Refrective index np	Viscosity (centipoises)
		(liquid)	(gas)		-				
Diborane	27,688	0.4818 <sup>-129.5</sup> (12)	-44 <sup>25</sup> (3)	3.1 <sup>18</sup> (15)	-92.5 (2)	-165.5 (2)	212.23 <sup>-112</sup> (16)	P4444444	************
		(liquid)	(gas)					(gas)	
Fluorine	38.000	1.14-200 (16)	<sub>0</sub> 25	1.60 <sup>-188</sup> (15)	-188 (15)	-833.0	760 <sup>-187</sup> •92	1.000195 <sup>20</sup>	
		(liquid)	(gas)						
Fluorine oxide	54,000	1.90 <sup>-223</sup> .8	5.5 <sup>25</sup> (14)	2.65 <b>-144.</b> 9 (14)	-144.9 (14)	-223.8 (2)	760 <sup>-144.9</sup> (14)		
		(liquid)	(gas)					(liquid)	(liquid)
Oxygen	32.000	1.14-183	<sub>0</sub> 25	1.629-183	-183.0	-218.4	760-182.97	1.221-181	0.189-252.07 (96%)
		(2)		(15)	(8)	(2)	(14)	(16)	(16)
		(liquid)	(liquid)					(liquid)	(liquid)
Hydrogen peroxide	34.016	1.436 <sup>25</sup> a(13)	-45.2 <sup>18</sup> (15)	11.61 <sup>18</sup> (15)	152.1	-0.89	2.1 <sup>25</sup> (14)	1.4139 <sup>22</sup> (13)	1.307 <sup>18</sup> (13)

aExtrapolated.

TABLE V - CALCULATED PERFORMANCE OF DIBORANE

			Mann	Froze	Frozen expansion			quilibrium e	xpansion	
(percent pell by dens weight) (gre	Pro- pellant density (gram/ cc)	Combustion- chamber temperature T <sub>C</sub> ( <sup>O</sup> K)	Mean molecular weight in combustion chamber, M <sub>C</sub> (gram/mole)	Specific impulse, I (lb-sec/lb)	Volume specific impulse, I <sub>d</sub> lb-sec cu ft (62.43)	Temper- ature at nozzle exit, Te (°K)	Specific impulse, I (lb-sec/lb)	Volume specific impulse, Id [ lb-sec cu ft (62.43)	nozzle exit, Te	Mean molecular weight at nozzle exit, Me (gram/ mole)
				1	3 <sub>2</sub> H <sub>6</sub> + F <sub>2</sub>					
9.85 alo.83 13.63 16.69 17.11 19.54 23.29	1.005 .993 .961 .928 .924 .900	5560 5574 5240 4900 4835 4467 4040	24.00 23.53 22.23 20.80 20.62 19.72 18.76	289.8 292.8 298.0 298.7 298.1 293.7 288.5	291.2 290.8 286.3 277.3 275.5 264.3 249.5	2655 2667 2640 2481 2451 2287 2110	315.5 317.4 322.4 318.4 317.9 312.9 307.1	317.0 315.2 309.8 295.6 293.7 281.6 265.6	4270 4298 4113 3650 3600 3589 3122	26.87 26.30 24.59 22.73 22.56 21.68 20.55
	<del></del>		<del></del>		32H6 + F2O	<b>'</b>	<b></b>		· · · · · ·	
12.03 #14.60 20.41 22.51 25.48	1.403 1.329 1.187 1.143 1.086	4750 4701 4460 4309 4007	25.03 21.97 19.83 19.22 18.49	280.0 286.0 295.0 295.3 293.5	392.8 380.1 350.2 337.4 318.7	2432 2440 2373 2315 2216	302.3 509.2 516.2 514.7 511.5	424.2 410.9 375.3 359.6 338.0	3757 3749 3493 3357 3172	25.18 23.95 21.53 20.93 20.14
				;	B <sub>2</sub> H <sub>6</sub> + O <sub>2</sub>					
20.61 22.38 25.71 32.90 36.58 38.82	0.890 .873 .844 .786 .760 .745	3953 3990 4022 3915 3740 3500	23.98 23.10 21.50 18.42 17.11 16.47	261.3 267.5 278.3 295.4 299.9 294.7	232.4 233.6 234.8 232.3 227.9 219.5	2438 2462 8475 2580 2240 2086	270.9 278.2 288.2 305.3 311.4 303.5	241.0 242.8 243.2 240.1 236.7 226.1	3139 3170 3176 2980 2830 2481	26.08 25.16 23.38 19.78 18.21 17.27
				В	<sub>2</sub> н <sub>6</sub> + н <sub>2</sub> о <sub>2</sub>	-				
7.53 411.95 21.34 28.93 35.18 37.38	1.250 1.161 1.009 .913 .846 .825	2858 3218 3204 3020 2846 2473	22.48 21.41 17.99 15.69 14.16 13.60	250.6 251.2 271.7 281.2 285.8 270.2	288.2 291.7 274.2 256.7 241.9 223.0	1780 2042 1980 1814 1672 1422	254.9 259.1 277.4 284.3 289.9 279.2	293.6 300.9 280.0 259.6 245.4 230.4	1968 2467 2197 1951 1899 1845	22.95 22.56 18.50 15.95 14.69 14.80

<sup>a</sup>Stoichiometric mixture

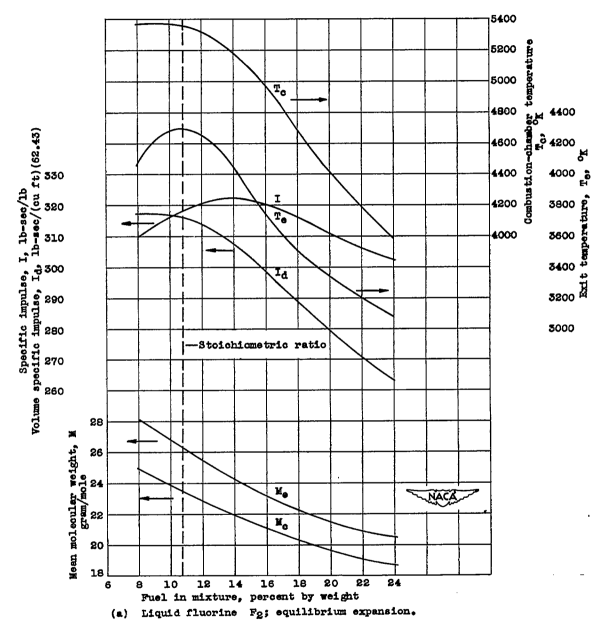


Figure 1. - Theoretical performance of diborane with liquid fluorine, liquid fluorine oxide, liquid oxygen, and 100-percent hydrogen peroxide. Isentropic expansion from 20.4 to 1 atmosphere.





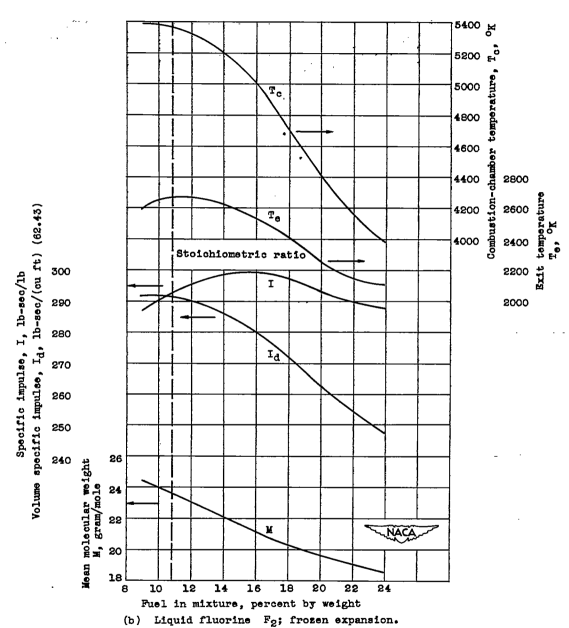


Figure 1. - Continued. Theoretical performance of diborane with liquid fluorine, liquid fluorine oxide, liquid oxygen, and 100-percent hydrogen peroxide. Isentropic expansion from 20.4 to 1 atmosphere.



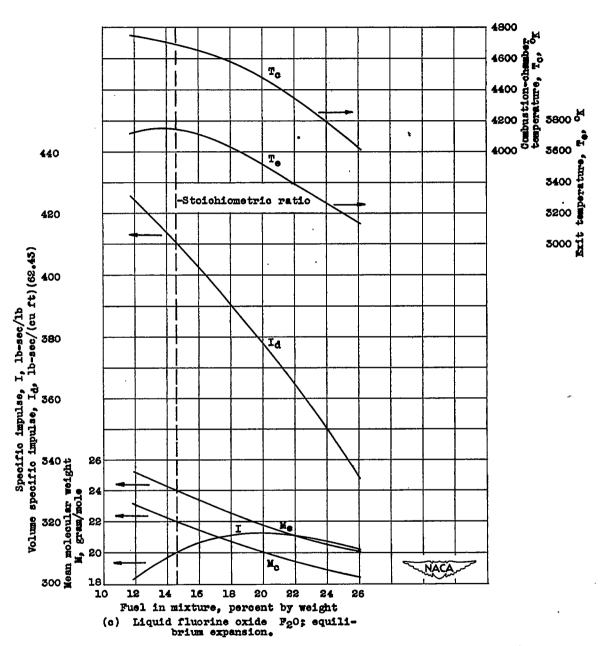
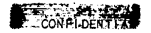


Figure 1. - Continued. Theoretical performance of diborane with liquid fluorine, liquid fluorine oxide, liquid oxygen, and 100-percent hydrogen peroxide. Isentropic expansion from 20.4 to 1 atmosphere.





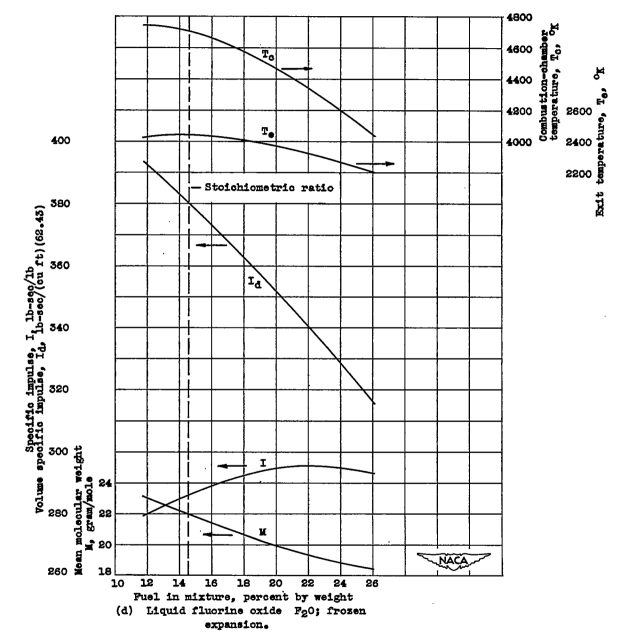
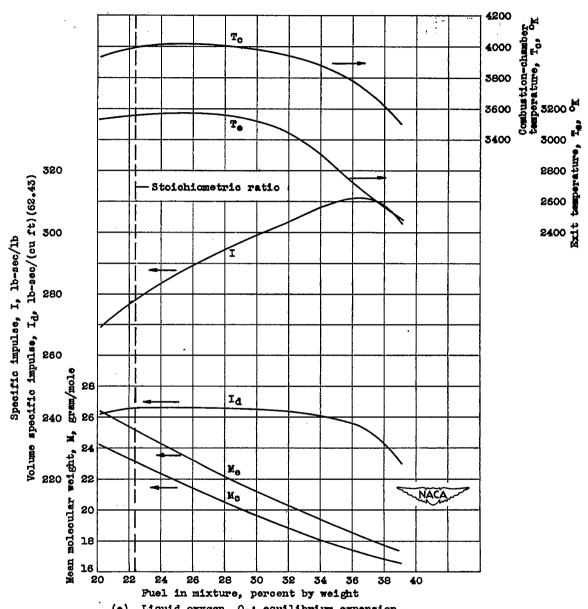


Figure 1. - Continued. Theoretical performance of diborane with liquid fluorine, liquid fluorine oxide, liquid oxygen, and 100-percent hydrogen peroxide. Isentropic expansion from 20.4 to 1 atmosphere.



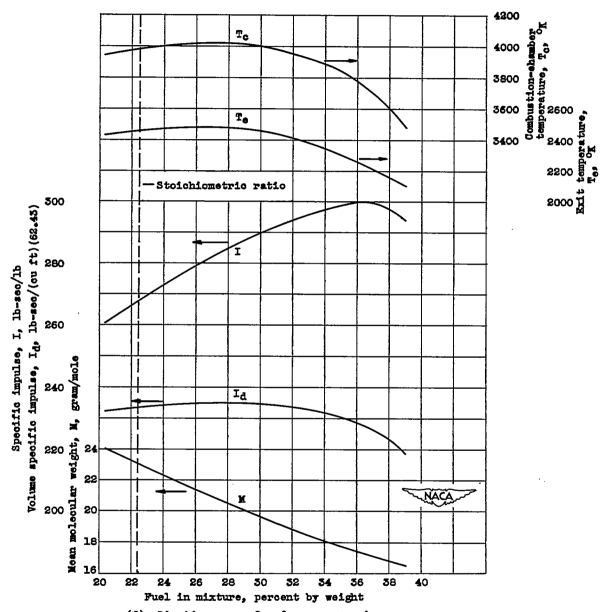


(e) Liquid oxygen O2; equilibrium expansion.

Figure 1. - Continued. Theoretical performance of diborane with liquid fluorine, liquid fluorine oxide, liquid oxygen, and loo-percent hydrogen peroxide. Isentropic expansion from 20.4 to 1 atmosphere.



TC2



(f) Liquid oxygen O2; frosen expansion.

Figure 1. - Continued. Theoretical performance of diborane with liquid fluorine, liquid fluorine oxide, liquid oxygen, and 100-percent hydrogen peroxide. Isentropic expansion from 20.4 to 1 atmosphere.

270

260

250

240

230 tg 20

220 Volume

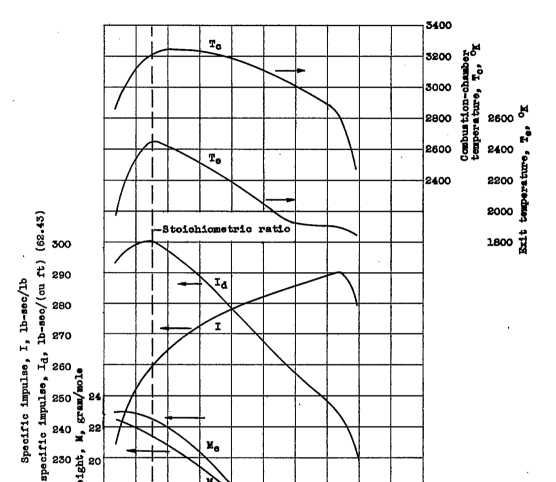
18

16

reinselom 14

9 12

10



Fuel in mixture, percent by weight (g) Hydrogen peroxide (100-percent) H<sub>2</sub>0<sub>2</sub>; equilibrium expansion. .

26

30

NACA

Figure 1. - Continued. Theoretical performance of diborane with liquid fluorine, liquid fluorine oxide, liquid oxygen, and lO0-percent hydrogen peroxide. Isentropic expansion from 20.4 to 1 atmosphere.

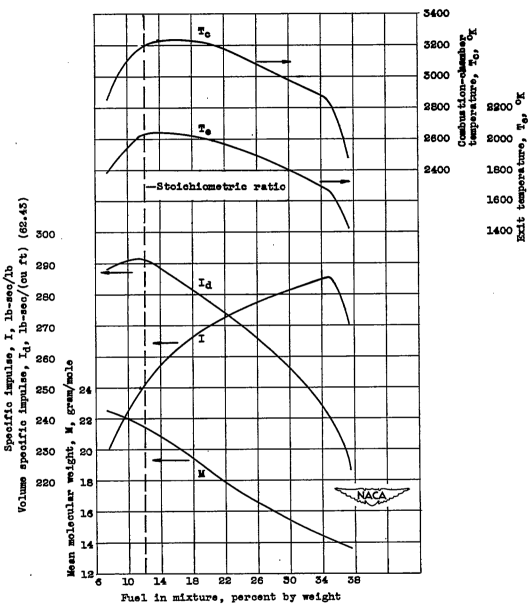
22

I

ĸe

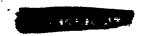
18

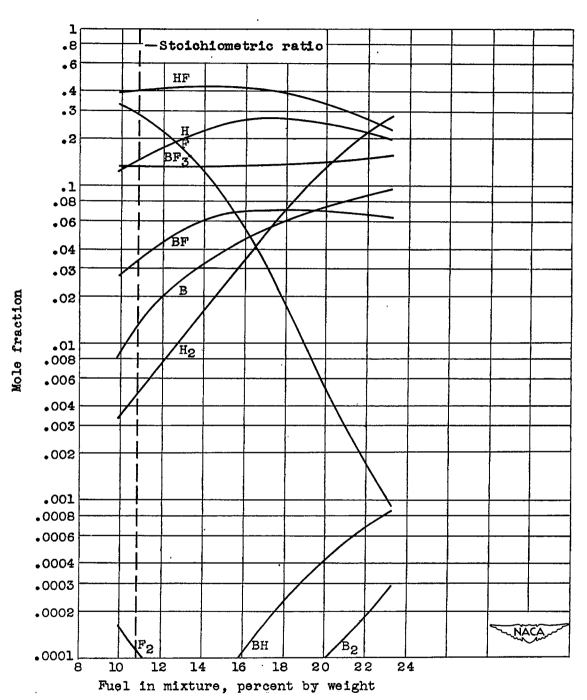




(h) Hydrogen peroxide (100-percent) H<sub>2</sub>0<sub>2</sub>; frozen expansion.

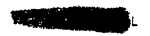
Figure 1. - Concluded. Theoretical performance of diborane with liquid fluorine, liquid fluorine oxide, liquid oxygen, and 100-percent hydrogen peroxide. Isentropic expansion from 20.4 to 1 atmosphere.



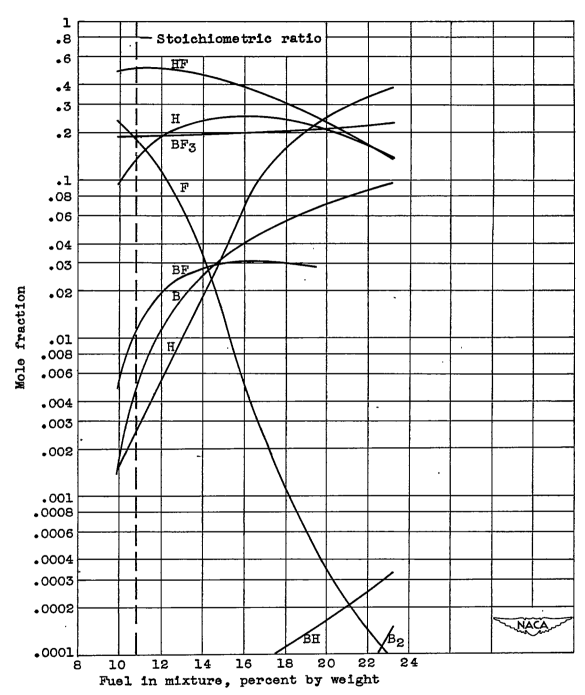


(a) Liquid fluorine F2; combustion-chamber conditions.

Figure 2. - Composition of products of reaction of diborane with liquid fluorine oxide, liquid oxygen, and 100-percent hydrogen peroxide.

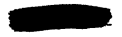


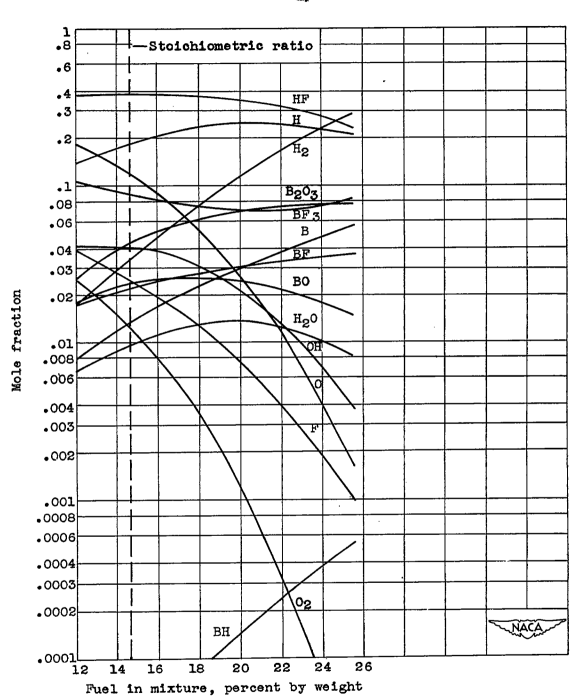




(b) Liquid fluorine F<sub>2</sub>; equilibrium nozzleexit conditions.

Figure 2. - Continued. Composition of products of reaction of diborane with liquid fluorine oxide, liquid oxygen, and loopercent hydrogen peroxide.

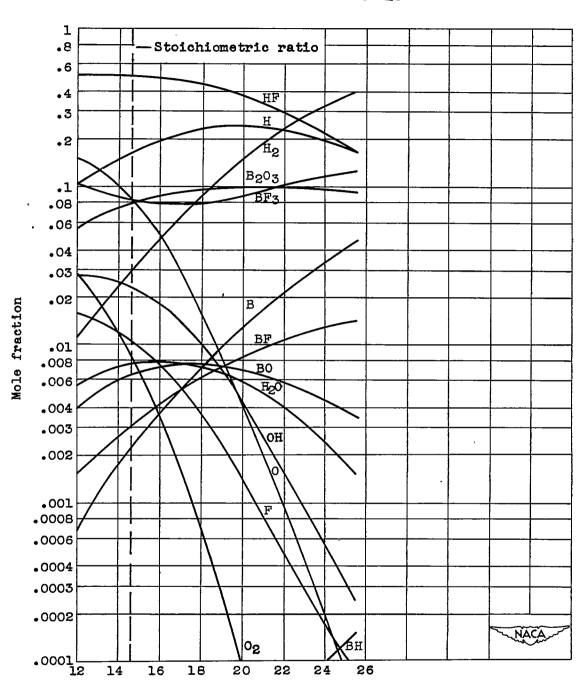




(c) Liquid fluorine oxide F20; combustion-chamber conditions.

Figure 2. - Continued. Composition of products of reaction of diborane with liquid fluorine oxide, liquid oxygen, and loopercent hydrogen peroxide.



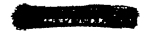


.....

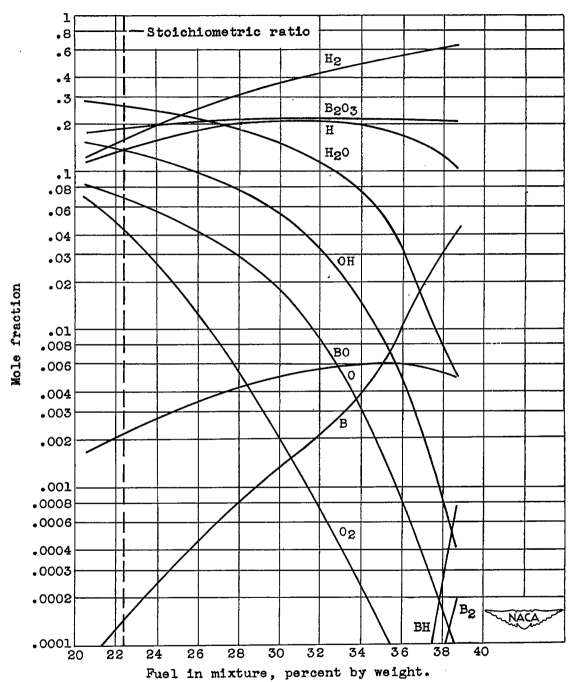
Fuel in mixture, percent by weight.

(d) Liquid fluorine oxide F<sub>2</sub>0; equilibrium nozzle-exit conditions.

Figure 2. - Continued. Composition of products of reaction of diborane with liquid fluorine oxide, liquid oxygen, and 100-percent hydrogen peroxide.

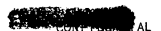


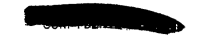


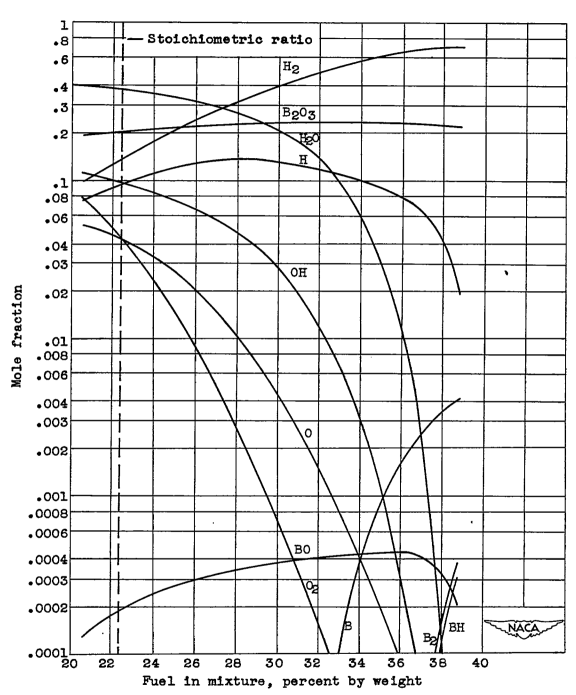


(e) Liquid oxygen 02; combustion-chamber conditions.

Figure 2. - Continued. Composition of products of reaction of diborane with liquid fluorine oxide, liquid oxygen, and loopercent hydrogen peroxide.

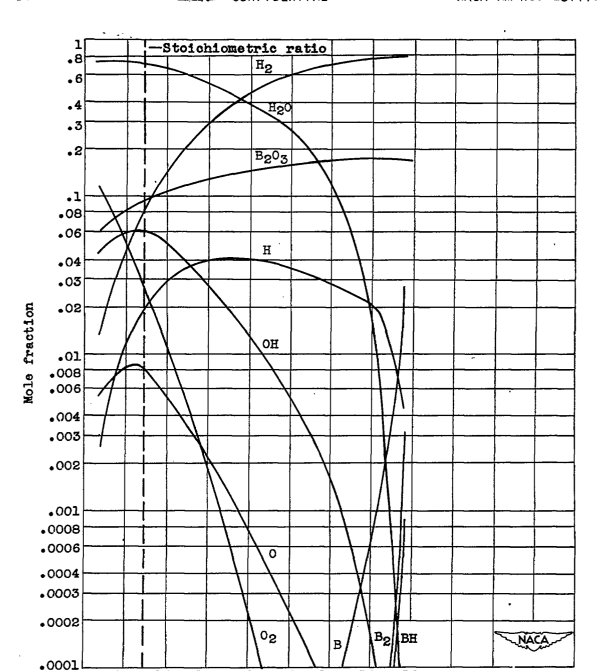






(f) Liquid oxygen 02; equilibrium nozzle-exit conditions.

Figure 2. - Continued. Composition of products of reaction of diborane with liquid fluorine oxide, liquid oxygen, and 100-percent hydrogen peroxide.



Fuel in mixture, percent by weight.

22

18

(g) Hydrogen peroxide (100 percent) H<sub>2</sub>0<sub>2</sub>; combustion-chamber conditions.

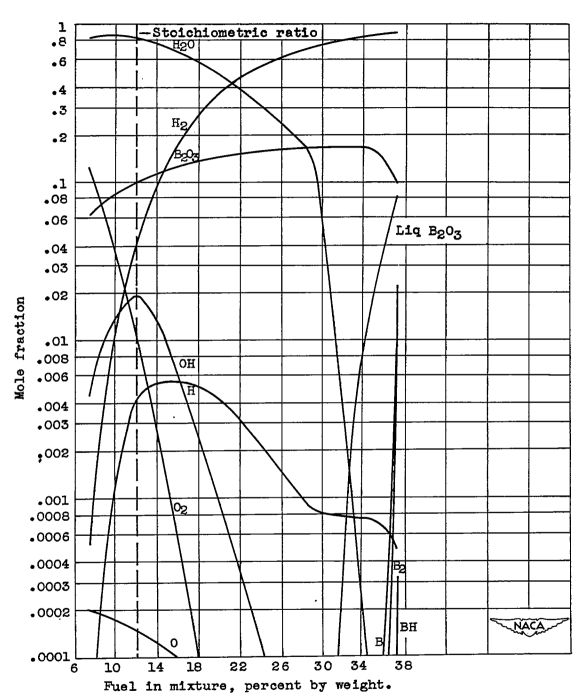
Figure 2. - Continued. Composition of products of reaction of diborane with liquid fluorine oxide, liquid oxygen, and 100-percent hydrogen peroxide.

26

30







(h) Hydrogen peroxide (100 percent) H<sub>2</sub>O<sub>2</sub>; equilibrium nozzle-exit conditions.

Figure 2. - Concluded. Composition of products of reaction of diborane with liquid fluorine oxide, liquid oxygen, and 100-percent hydrogen peroxide.

